=> s xylene

L1 93972 XYLENE

=> s ?decene

L2 15629 ?DECENE

=> s 11 and 12

L3 504 L1 AND L2

=> s sulfonic acid

L4 55946 SULFONIC ACID

=> s 14 and 13

L5 10 L4 AND L3

=> d 15 1-10 abs

L5

L5 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2002 ACS

AB A lubricant compn. comprising an oligomeric ketone which is a fluid and which is obtainable by reacting an alkene contg. from 4 to 20 carbon atoms with carbon monoxide in the presence of a catalyst which is obtainable by combining a transition metal or a compd. thereof and a bidentate ligand.

L5 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2002 ACS

- This invention relates to a solid acid catalyst comprising porous highly fluorinated acidic polymer and process for manufg. it. The catalyst comprises at least 90% highly fluorinated acidic polymer and has a surface area of .apprx.0.01 m2/g to .apprx.5 m2/g. The catalyst is prepd. by exposing the polymer to a pressurized fluid such as SO2 or CO2, heating the polymer to above the softening point, reducing the pressure quickly thus expanding the polymer into a porous structure, and cooling the polymer. Examples of the highly fluorinated acidic polymer are perfluorinated sulfonic acid resin perfluorinated and sulfonyl fluoride polymer.
- L5 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2002 ACS
- AB A new versatile reagent, a perfluorosulfonate/trisilanol [(OH)3Si(CH2)3(CF2)2O(CF2)2SO3-M+], is synthesized and a series of surface-bound strong solid acid catalysts developed which are active for catalyzing a range of reactions (e.g., alkene isomerization, alkylations, and acylations). This catalyst, when bound on SiO2, showed significantly higher activity for benzene and toluene alkylation, m-xylene benzoylation, and isomerization of 1-butene than Nafion NR5O and Amberlyst-15 catalysts.
- L5 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2002 ACS
- AB Control by best demonstrated technol. of air pollution by org. compds. emitted during the reaction of feedstocks or chem. intermediates other than air in reactors in the manuf. of 173 specific chems. is established under the Federal Clean Air Act. All new, modified, and reconstructed distn. facilities must reduce volatile org. compd. (total org. compds., minus CH4 and C2H6) emissions by 98% by wt. or to a concn. of 20 ppm by vol. on a dry basis cor. to 3% O, whichever is less stringent, or combust the emissions in a flare, or maintain a TRE (total resource effectiveness) index value >1.0 without the use of emission control devices. Methods of reducing emissions may include incineration, introduction of the vent stream into the flame zone of a boiler or process heater, or recovery with absorbers, condensers, or other devices. Reactor facilities with flow rates .ltoreq.0.011 scm/min, facilities in process units with prodn. capacities .ltoreq.1 Gg/yr, facilities with TRE index values >8.0, and batch processes are exempt from all or part of the std.

Control by best demonstrated technol. of air pollution by org. compds. AΒ emitted during distn. processes in the prodn. of 211 specific chems. is established under the Federal Clean Air Act. All new, modified, and reconstructed distn. facilities must reduce volatile org. compd. (total org. compds. minus CH4 and C2H6) emissions by 98% by wt. or to a concn. of 20 ppm by vol. on a dry basis cor. to 3% O, whichever is less stringent, or combust the emissions in a flare, or maintain a TRE (total resource effectiveness) index value >1.0 without the use of emission control devices. Methods of reducing emissions may include incineration, introduction of the vent stream into the flame zone of a boiler or process heater, or recovery with absorbers, adsorbers, condensers, or other devices. Distn. facilities that are a part of a unit that produces coal tar or beverage alc., that are designed and operated as a batch operation, that are a part of a polymer manufg. process subject to the polymer manuf. new source performance stds., or that use, contain, and produce no volatile org. compds. are exempt. Distn. facilities with TRE index values >8.0 are exempt from a portion of the requirements.

ANSWER 6 OF 10 CAPLUS COPYRIGHT 2002 ACS

The title resins contain .gtoreq.1 linkage of -[Si(R1)(R2)O]mMR3nR41 (M = Cu, Mg, Sr, Ba, Zn, Sn, Pb, Zr, Mn, Fe, Co, Ni; R1 = H, OH, alkoxy, C1-18 alkyl, aryl; R2 = H, OH, alkoxy, C1-18 alkyl groups, aryl, siloxane, polysiloxane; R3 = OH, alkoxy, modifier residue groups; R4 = nonreactive O, H, alkoxy; m = .gtoreq. 1 integer; n = 0, >1 integer; 1 = 0-2; such that the sum of m + n + 1 is equal to the valence of M, and the R1 and R3 optionally form rings via metals having valence >2). Xylene 50, Me2Si(OMe)2 50, Cu hydroxide 6, and octanol 1.8 parts were heated at 70-110.degree. with removal of MeOH to prep. a varnish.

- ANSWER 7 OF 10 CAPLUS COPYRIGHT 2002 ACS

  Desired product distributions of alkylarom. hydrocarbons are obtained by choosing appropriate reaction conditions for Friedel-Crafts alkylation of .gtoreq.1 arom. hydrocarbon with .gtoreq.1 .alpha.-alkene. Suitable reactants include C8-24-.alpha.-olefins, C6H6, PhMe, isopropylbenzene, xylene, ethylbenzene, and tri- and tetramethylbenzenes. The alkylarom. hydrocarbons are then sulfonated to produce surfactants suitable for use in petroleum recovery. PhMe was alkylated with 1-dodecene in the presence of AlCl3 at 7:1 PhMe-dodecene mol ratio and at 20.degree. intervals from -20.degree. to 60.degree. The relative distribution of C2-vs. C4-alkylated products (in mol%) were 67.9:12.9 at -20.degree., 68.2:3.1 at 0.degree., 60.8 = 18.5 at 20.degree., 39.6:40.4 at 40.degree., and 41.2:37.85 at 60.degree..
- ANSWER 8 OF 10 CAPLUS COPYRIGHT 2002 ACS

  Monoalkyl sulfates were prepd. by reaction of hydrocarbon oils contg. olefinic, arom., or satd. aliph. hydrocarbons with 70-90 wt.% H2SO4 at <100.degree. Thus, a hydrocarbon oil contg. 1-dodecene 20, xylene 35, and dodecane 45 wt.% was treated with 80 wt.% H2SO4 at 30.degree. for 4 h to give 95% esterification and 5% alkylation with 75% conversion, vs. 5, 95, and 93%, resp., with 95 wt.% H2SO4.
- L5 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2002 ACS

  AB Two new methods are presented for prepg. HO sulfonic

  acids (I) as well as an improved procedure for dehydrating I to
  sultones. 3-Hydroxy-1-nonadecene is treated with NaHSO3 and a
  peroxide to give 83% Na 3-hydroxy-1-nonadecane-sulfonate. The
  corresponding acid, when refluxed in xylene gives the 59%
  sultone. Adding NaHSO3 to an .alpha.,.beta.-unsatd. ketone and reducing
  with Raney Ni gives I. E.g. mesityl oxide gives Na 2-methyl-4-hydroxy-2pentanesulfonate; refluxing the free acid in xylene gives 70%
  sultone.
- L5 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2002 ACS
  AB High-mol. alcs. are prepd. by heating satd. primary aliphatic alcs. contg.

more than 8 C atoms/mol. at temps. above 100.degree. in the presence of minor quantities of org. sulfonic acids and sometimes of solvents or diluents. The resultant alcs., contg. twice as many C atoms/mol. as the initial material, are useful as intermediates for the prepn. of auxiliary agents for textiles. Heating dodecyl alc. 200 and p-MeC6H4SO3H (I) 10 parts 6 hrs. at 220-50.degree. in a N atm. yields a residue which on vacuum distn. gives tetracosanol 142, b7.5-9 221-46.degree., as well as an oil 48, bl0 85-191.degree., consisting of a mixt. of dodecene and dimerized products. Analogously, octadecanol, I, and xylene give a product, m. 59-9.5.degree. (from C6H6), mol. wt. 536, OH index 93, probably hexatriacontanol.

=> d his

(FILE 'HOME' ENTERED AT 22:47:28 ON 20 MAR 2002)

FILE 'CAPLUS, CAOLD' ENTERED AT 22:47:50 ON 20 MAR 2002
L1 93972 S XYLENE
L2 15629 S ?DECENE

L3 504 S L1 AND L2 L4 55946 S SULFONIC ACID L5 10 S L4 AND L3

=> d 15 7-8 ibib kwic

L5 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1987:620278 CAPLUS

DOCUMENT NUMBER: 107:220278

TITLE: Selective alkylation of aromatic hydrocarbons

INVENTOR(S): Aldrich, Haven S.; Puerto, Maura C.; Reed, Ronald L.

PATENT ASSIGNEE(S): Exxon Production Research Co., USA

SOURCE: Fr. Demande, 18 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

FR 2589858 A1 19870515 FR 1986-11522 19860808
PRIORITY APPLN. INFO.: US 1985-798074 19851114

Desired product distributions of alkylarom. hydrocarbons are obtained by choosing appropriate reaction conditions for Friedel-Crafts alkylation of .gtoreq.1 arom. hydrocarbon with .gtoreq.1 .alpha.-alkene. Suitable reactants include C8-24-.alpha.-olefins, C6H6, PhMe, isopropylbenzene, xylene, ethylbenzene, and tri- and tetramethylbenzenes. The alkylarom. hydrocarbons are then sulfonated to produce surfactants suitable for use in petroleum recovery. PhMe was alkylated with 1-dodecene in the presence of AlCl3 at 7:1 PhMe-dodecene mol ratio and at 20.degree. intervals from -20.degree. to 60.degree.. The relative distribution of C2-vs. C4-alkylated products (in mol%) were 67.9:12.9 at -20.degree., 68.2:3.1 at 0.degree., 60.8 = 18.5 at 20.degree., 39.6:40.4 at 40.degree., and 41.2:37.85 at 60.degree..

IT Aromatic hydrocarbons, reactions

RL: RCT (Reactant)

(C6-8, alkylation of, with dodecene, product distribution in)

IT Sulfonic acids, preparation

RL: PREP (Preparation)

(alkylarene, prepn. of, Friedel-Crafts alkylation in, product distribution of)

IT 112-41-4, 1-Dodecene

RL: USES (Uses)

(alkylation of alkylarom. hydrocarbons with, product distribution in) 98-82-8, Isopropylbenzene 71-43-2, Benzene, reactions ΙT Ethylbenzene, reactions 1330-20-7, Xylene, reactions 25619-60-7, Tetramethylbenzene 25551-13-7, Trimethylbenzene RL: RCT (Reactant) (alkylation of, with alkenes, product distribution in) ΙT 108-88-3, Toluene, reactions RL: RCT (Reactant)

(alkylation of, with dodecene, product distribution in)

ANSWER 8 OF 10 CAPLUS COPYRIGHT 2002 ACS 1982:162124 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

96:162124

TITLE:

SOURCE:

Monoalkyl sulfates

PATENT ASSIGNEE(S):

Nippon Mining Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
		'		
JP 56164162	A2	19811217	JP 1980-65895	19800520
JP 63052624	B4	19881019		

Monoalkyl sulfates were prepd. by reaction of hydrocarbon oils contg. AB olefinic, arom., or satd. aliph. hydrocarbons with 70-90 wt.% H2SO4 at <100.degree.. Thus, a hydrocarbon oil contg. 1-dodecene 20, xylene 35, and dodecane 45 wt.% was treated with 80 wt.% H2SO4 at 30.degree. for 4 h to give 95% esterification and 5% alkylation with 75%conversion, vs. 5, 95, and 93%, resp., with 95 wt.% H2SO4. ΙT Esterification

(of aliph. hydrocarbons, with sulfonic acid, monoalkyl sulfates from)

```
=> d his
     (FILE 'HOME' ENTERED AT 22:47:28 ON 20 MAR 2002)
     FILE 'CAPLUS, CAOLD' ENTERED AT 22:47:50-ON 20 MAR 2002
          93972 S XYLENE
L1
          15629 S ?DECENE
L2
L3
            504 S L1 AND L2
          55946 S SULFONIC ACID
L4
L5
             10 S L4 AND L3
              0 S DODECECE
L6
           3415 S DODECENE
L7
            160 S L7 AND L1
L8
              O S L8 AND (METHANE SULFONIC ACID)
L9
            330 S METHANE SULFONIC ACID
L10
              1 S METHANE SULPHONIC ACID
L11
     FILE 'CAPLUS, CAOLD' ENTERED AT 23:12:57 ON 20 MAR 2002
            330 S METHANE SULFONIC ACID
L12
              0 S L12 AND L7
L13
             11 S L12 AND L1
L14
L15
            116 S ALKANE SULFONIC ACID
L16
              0 S L15 AND L1 AND L2
              2 S L12 AND L2
L17
L18
             46 S METHYL SULFONIC ACID
              O S (METHYL SULFONIC ACID)/RN
L19
     FILE 'REGISTRY' ENTERED AT 23:58:44 ON 20 MAR 2002
              0 S (METHYL SULFONIC ACID)/RN
L20
L21
              O S (METHANE SULFONIC ACID)/RN
L22
              O S (METHANE SULFONIC ACID)/CN
              O S (METHANE SULFONIC ACID)/CRN
L23
              0 S ACTIC ACID/RN
L24
L25
              O S (METHANE SULFONIC ACID)/CRN
     FILE 'REGISTRY' ENTERED AT 00:01:33 ON 21 MAR 2002
L26
              O S (METHANE SULFONIC ACID)/CN
L27
              1 S (ACETIC ACID)/CN
              O S (METHANE SULPHONIC ACID)/CN
L28
              O S (METHYL SULPHONIC ACID)/CN
L29
                STRUCTURE UPLOADED
L30
L31
             47 S L30
L32
              1 S (75-75-2)/RN
     FILE 'CAPLUS, USPATFULL, CASREACT' ENTERED AT 00:08:49 ON 21 MAR 2002
     FILE 'CAPLUS, CAOLD, USPATFULL' ENTERED AT 00:08:59 ON 21 MAR 2002
           4156 S (75-75-2)/RN
L33
L34
            153 S L33 AND ALKYLATION
L35
              0 S L34 AND OLEFINE
L36
             50 S L34 AND OLEFIN
L37
             33 S L36 AND (XYLENE OR BENZENE OR TOLUENE)
L38
           4162 S 37 AND DECENE
              7 S L37 AND DECENE
```

=>

L39

```
=> d his
L1
```

(FILE 'HOME' ENTERED AT 22:47:28 ON 20 MAR 2002)

```
FILE 'CAPLUS, CAOLD' ENTERED AT 22:47:50 ON 20 MAR 2002
          93972 S XYLENE
L2
          15629 S ?DECENE
            504 S L1 AND L2
L3
          55946 S SULFONIC ACID
L4
L5
             10 S L4 AND L3
              0 S DODECECE
L6
           3415 S DODECENE
L7
            160 S L7 AND L1
^{18}
L9
              O S L8 AND (METHANE SULFONIC ACID)
            330 S METHANE SULFONIC ACID
L10
              1 S METHANE SULPHONIC ACID
L11
     FILE 'CAPLUS, CAOLD' ENTERED AT 23:12:57 ON 20 MAR 2002
L12
            330 S METHANE SULFONIC ACID
              0 S L12 AND L7
L13
             11 S L12 AND L1
L14
L15
            116 S ALKANE SULFONIC ACID
L16
              0 S L15 AND L1 AND L2
=> s 112 and 12
             2 L12 AND L2
L17
=> d 117 1-2 ibib kwic
```

L17 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS 2001:168087 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:210386

TITLE: Lubricants containing oligomeric polyketones

Boyde, Stephen; Cameron, Paul Alexander; Eastham, INVENTOR(S):

Graham Ronald; Tooze, Robert Paul

PATENT ASSIGNEE(S): Imperial Chemical Industries PLC, UK

SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

```
PATENT NO. KIND DATE
                                                     APPLICATION NO. DATE
                                                WO 2000-GB3235 20000822
                            ----
      WO 2001016260
                           A1 20010308
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
                CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
                HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
                YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
           RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                    GB 1999-20516
                                                                          A 19990901
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
                                MARPAT 134:210386
REFERENCE COUNT:
                                2
                                        THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS
                                        RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                            75-75-2, Methane sulfonic
      67-56-1, Methanol, reactions
```

TT 91-13-4, .alpha.,.alpha.'-Dibromo-o-xylene 106-51-4, 1,4-Benzoquinone, reactions 109-72-8, Butyl lithium, reactions 630-08-0, Carbon monoxide, reactions 829-85-6, Diphenylphosphine 872-05-9, 1-Decene 3375-31-3 6131-90-4, Sodium acetate 67969-84-0, trihydrate 6737-42-4, 1,3-Bis(diphenylphosphino)propane 1,3-Dibromo-2,2-diethyl propane 328952-85-8 RL: RCT (Reactant); RACT (Reactant or reagent) (lubricants contg. oligomeric polyketones)

L17 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1989:616732 CAPLUS

111:216732 DOCUMENT NUMBER:

Recovery of transition metals from aqueous solutions TITLE: Goodall, Brian Leslie; Grotenhuis, Paulus Alexander INVENTOR(S):

Maria

PATENT ASSIGNEE(S): Shell Internationale Research Maatschappij B. V.,

Neth.

Brit. UK Pat. Appl., 16 pp. SOURCE:

CODEN: BAXXDU

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

١

GB 2213907 APPLICATION NO. DATE \_\_\_\_\_ GB 2213807 A1 19890823 US 4880546 A 19891114 GB 1987-29446 19871217 US 1988-285464 19881216 19890823 PRIORITY APPLN. INFO.: GB 19
OTHER SOURCE(S): CASREACT 111:216732 GB 1987-29446 19871217

872-05-9, 1-Decene

RL: RCT (Reactant)

(hydroformylation of, spent catalyst from, rhodium recovery from)

ΙT 75-75-2, Methane sulfonic acid

RL: USES (Uses)

(in ethene carbonylation with methanol, palladium-contg. catalyst for)

## WEST

Generate Collection Print

L1: Entry 1 of 3

File: EPAB

May 15, 1987

PUB-NO: FR002589858A1

DOCUMENT-IDENTIFIER: FR 2589858 A1

TITLE: Process for selective alkylation of aromatic compounds

PUBN-DATE: May 15, 1987

INVENTOR-INFORMATION:

NAME COUNTRY

ALDRICH, HAVEN S PUERTO, MAURA C REED, RONALD L

ASSIGNEE-INFORMATION:

NAME

EXXON PRODUCTION RESEARCH CO US

APPL-NO: FR08611522

APPL-DATE: August 8, 1986

PRIORITY-DATA: US79807485A (November 14, 1985)

US-CL-CURRENT:  $\frac{585}{447}$  INT-CL (IPC):  $\frac{507C}{2/66}$ 

EUR-CL (EPC): C07C002/66; E21B043/22

ABSTRACT:

Process for preparing a reaction product having a desired distribution.

This product is obtained by carrying out a Friedel-Crafts alkylation reaction of one or more aromatic compounds with one or more linear alpha olefins at a preferred reaction temperature and sulphonation of the reaction product.

Application to the preparation of a surfactant used in the assisted recovery of oil.

## WEST

Generate Collection Print

L1: Entry 2 of 3

File: DWPI

May 15, 1987

DERWENT-ACC-NO: 1987-171983

DERWENT-WEEK: 198725

COPYRIGHT 2002 DERWENT INFORMATION LTD

TITLE: Obtaining desired isomer distribution from friedel-crafts alkylation - of aromatic cpd. with linear alpha-olefin by adjusting reaction temp., surfactant for enhanced oil recovery then being obtd. by sulphonation

INVENTOR: ALDRICH, H S; PUERTO, M C ; REED, R L

PATENT-ASSIGNEE:

ASSIGNEE CODE EXXON PRODN RES CO ESSO

PRIORITY-DATA: 1985US-0798074 (November 14, 1985)

PATENT-FAMILY:

PUB-NO PUB-DATE LANGUAGE PAGES MAIN-IPC

FR 2589858 A May 15, 1987 018

APPLICATION-DATA:

PUB-NO APPL-DATE APPL-NO DESCRIPTOR

FR 2589858A August 8, 1986 1986FR-0011522

INT-CL (IPC): C07C 2/68; C07C 15/10; C07C 143/34; C09K 7/02; E21B 43/22

ABSTRACTED-PUB-NO: FR 2589858A

BASIC-ABSTRACT:

Process for obtaining the desired distribution of a reaction prod. from a Friedel-Crafts alkylation reaction of aromatic cpd.(s) with linear alpha-olefin(s) consists of: (a) determining experimentally the distribution of the reaction prod. at various reaction temps.; (b) from the results of (a), choosing a pref. reaction temp., enabling the desired distribution to be obtd.; and (c) carrying out the reaction at the pref. temp.

The aromatic cpd. may be benzene, toluene, isopropylbenzene, xylene, ethylbenzene, tri-tetramethylbenzene or their mixts. The linear alpha-olefins may be 8-24C.

USE/ADVANTAGE - With the further step of sulphonating the reaction prod., surfactants can be produced for enhanced oil recovery, with isomer distribution adjusted to the conditions of the reservoir. It may be then be possible to omit the alcohol previously added to micellar solns. of alkylaryl sulphonates. /2

TITLE-TERMS: OBTAIN ISOMER DISTRIBUTE FRIEDEL CRAFT ALKYLATED AROMATIC COMPOUND LINEAR ALPHA OLEFIN ADJUST REACT TEMPERATURE SURFACTANT ENHANCE OIL RECOVER OBTAIN SULPHONATED

DERWENT-CLASS: E14 H01 H04 Q49

CPI-CODES: E10-J02B3; H01-D06; H04-E; N04-D01;

## CHEMICAL-CODES:

Chemical Indexing M3 \*01\*
 Fragmentation Code
 G010 G011 G012 G013 G014 G015 G016 G017 G018 G020
 G021 G022 G029 G040 G100 M210 M211 M212 M213 M220
 M222 M223 M224 M225 M226 M231 M232 M240 M281 M282
 M283 M320 M414 M510 M520 M531 M540 M720 M903 M904
 N203 N213 N221 N311 N318 N441 N511 N512 N513
 Markush Compounds
 198725-B3201-P
 Registry Numbers
 87140 1286M

Chemical Indexing M3 \*02\*
Fragmentation Code
A313 A940 C017 C100 C730 C801 C803 C804 C805 C806
C807 M411 M730 M903 M910 Q421
Registry Numbers
87140 1286M

## UNLINKED-DERWENT-REGISTRY-NUMBERS: 0862S; 1677S

SECONDARY-ACC-NO:

CPI Secondary Accession Numbers: C1987-071644 Non-CPI Secondary Accession Numbers: N1987-129062